PATENT COOPERATION TREATY

PCT

INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY

(Chapter II of the Patent Cooperation Treaty)

(PCT Article 36 and Rule 70)

REC'D 3 1 MAR 2006

				VVIPO FOI
• •	cant's or agent's file reference 003/254/PCT	FOR FURTHER AC	TION	See Form PCT/IPEA/416
International application No. PCT/CU2004/000012		International filing date (day/month/year)	Priority date (day/month/year) 04.11.2003
	ational Patent Classification C07K14/22 A61K39/	on (IPC) or national classification and IF		
Applic		IA GENETICA Y BIOTECNOLO	GIA	
1.	This report is the inter Authority under Article	national preliminary examination re 35 and transmitted to the applican	port, established by the according to Article 3	nis International Preliminary Examining 36.
2.	This REPORT consist	ts of a total of 7 sheets, including t	nis cover sheet.	
3.	This report is also acc	companied by ANNEXES, comprisi	ng:	
a. sent to the applicant and to the International Bureau) a total of 2 sheets, as				s, as follows:
	and/or she	the description, claims and/or drawing the containing rectifications authoritative Instructions).	ngs which have been a zed by this Authority (amended and are the basis of this report see Rule 70.16 and Section 607 of the
	☐ sheets wh beyond th Suppleme	e disclosure in the international app	hich this Authority con dication as filed, as inc	nsiders contain an amendment that goes dicated in item 4 of Box No. I and the
	egauanca listin	ternational Bureau only) a total of (ing and/or tables related thereto, in equence Listing (see Section 802 of	celectronic torm only, a	ber of electronic carrier(s)) , containing a as indicated in the Supplemental Box structions).
4.	This report contains i	ndications relating to the following i	tems:	
	⊠ Box No. I Bas	sis of the report		
		ority		
		n-establishment of opinion with reg	ard to novelty, inventiv	e step and industrial applicability
		ck of unity of invention		
	⊠ Box No V Re	asoned statement under Article 350 plicability; citations and explanation	2) with regard to nove supporting such stat	elty, inventive step or industrial sement
	•	rtain documents cited		
		rtain defects in the international ap	olication	
		rtain observations on the internatio		
			Data of completion of	this report
Date of submission of the demand			Date of completion of	uno report
20.	04.2005		30.03.2006	
Nan prei	ne and mailing address of liminary examining author	ity:	Authorized officer	Sportternes Peterson, Fig.
_	NL-2280 HV R Tel. +31 70 34	ent Office - P.B. 5818 Patentiaan 2 ijswijk - Pays Bas 0 - 2040 Tx: 31 651 epo ni	Hix, R	70 240 2808
. —	Fax: +31 70 34	HI + 2010	Telephone No. +31 7	し つせつ つつ つつ しつ しゅう は は で で で で で で で で で で で で で で で で で

INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY

International application No. PCT/CU2004/000012

Box No. I Basis of the report With regard to the language, this filed, unless otherwise indicated	report is based on the international application in the language in which it was			
With regard to the language, this filed, unless otherwise indicated	report is based on the international application in the language in which it was			
With regard to the language, this report is based on the international application in the language in which it was filed, unless otherwise indicated under this item.				
which is the language of a tr international search (und publication of the internation	ional application (under Rule 12.4)			
☐ international preliminary	examination (under Rules 55.2 and/or 55.3)			
Nith regard to the elements * of the international application, this report is based on <i>(replacement sheets which</i> have been furnished to the receiving Office in response to an invitation under Article 14 are referred to in this report as "originally filed" and are not annexed to this report):				
Description, Pages				
1-20	as originally filed			
Claims, Numbers				
1-12	received on 20.04.2005 with letter of 18.04.2005			
Drawings, Figures	\cdot			
1-11	as originally filed			
☐ a sequence listing and/or ar	y related table(s) - see Supplemental Box Relating to Sequence Listing			
 ☐ the description, pages ☐ the claims, Nos. ☐ the drawings, sheets/figs ☐ the sequence listing (sp. 	s ecify):			
had not been made, since they Supplemental Box (Rule 70.2(c)) the description, pages the claims, Nos. the drawings, sheets/figsthe the sequence listing (sp. any table(s) related to sp.	3			
	which is the language of a traditional search (under publication of the international preliminary of international preliminary of the elements* of thave been furnished to the receiverport as "originally filed" and are preparted as "originally fil			

INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY

Box No. V Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement

1. Statement

Novelty (N)

Yes: Claims

1-12

No: Claims

No:

Inventive step (IS)

Yes: Claims

2-4, 6-12

No: Claims

1,5

1-12

Industrial applicability (IA)

Yes: Claims

Claims

2. Citations and explanations (Rule 70.7):

see separate sheet

Box No. VII Certain defects in the international application

The following defects in the form or contents of the international application have been noted:

see separate sheet

- 1 The following documents are referred to;
- D1: EP-A-1 297 844 (Microbiological Research Authority)
- D2: Biochemical and Biophysical Research Communications, G. Sardiñas et al. vol. 277, pages 51-54 (2000)
- D3: US-A-2003/0059444 (Zollinger et al.)
- D4: Infection and Immunity, Saunders et al. Jan. 1999, pg. 113-119, vol. 67. no. 1
- D6: WO-A-03 051 379 (Microbiological Research Authority)
- D7: WO-A-01 91 788(Statens Insititutt for Folkehelse)
- D8: WO-A-01 09 350 (Smithkline Beechan Biologicals S.A.)
- D9: Infection and Immunity, Jin et al., vol. 71, no. 9, pages 5115-5120, Sept. 2003.

Re Item V

Reasoned statement with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement

- 2 **NOVELTY** (Art. 33(2) PCT)
- 2.1 D1 discloses compositions comprising *N.meningitidis* outer membrane vesicles (OMV) enriched with antigenic components used in the form of a vaccine in order to elicit an immune response. The antigens used in the present case are the Transferrin binding proteins TbpA and TbpB. The enrichment is achieved by mixing the OMV with the antigens.
- In view of the prior art cited, claims 1 to 12 appear to be novel and meet therefore the requirements of Art. 33(2) PCT, since the prior art does not disclose vaccines in which the antigen has been incorporated into the bacterial outer membrane vesicle by co-folding such that the vesicle structure is maintained intact.
- 3 **INVENTIVE STEP** (Art. 33(3) PCT)
- 3.1 Documents D3 and D4 are considered to represent the most relevant state of the

art and disclose a vaccine using native outer membrane vesicles (NOMV) for Neisseria or other Gram negative bacteria. D3, Page 1, paragraph 8, states that "the antigens presented as part of the NOMV... are in a completely native configuration and environment as part of intact outer membrane".

- The difference between the subject-matter of the present application and that disclosed in the prior art is that the prior art methods involve the generation of OMVs from genetically modified bacteria, compared to the present application which involves the incorporation of the antigens into the OMV.
- The problem to be solved by the present invention may therefore be regarded as providing a method of incorporation of protein antigens into outer membrane vesicles.
- The proposed solution is the incorporation of an antigen into a bacterial outer membrane vesicle using the method of claim 1 such that the vesicle structure is maintained intact and proper folding of the antigen is achieved.
- D2 describes the conjugation of the P64k peptide from *N.meningitidis* strain CU385 (B:4:P1.19, 15) outer membrane vesicles (OMV; as used in the present application) used as a carrier to two cyclic synthetic peptides derived from variable regions of the outer membrane protein PorA. The P64k was found to be an efficient carrier protein for PorA derived peptides. The chemical conjugation to the carrier did not affect the folding and allowed the synthetic peptides to induce a PorA-specific immune response.
- 3.5.1 Although the person skilled in the art is aware of the use of OMV as an effective carrier for antigens, resulting in an effective induction of an immune response, D2, the antigen was chemically conjugated to the OMV for use as a carrier in the prior art disclosure compared to the present application where the antigen is incorporated into the OMV by co-folding.
- 3.5.2 The use of outer membrane vesicles from gram negative bacteria is commonly

known in the state of the art for use as vaccines, see D6 to D8, to name but a few, however in the present application the OMV are not used as carriers but are used to refold the antigens which are incorporated into the OMV structure.

- 3.5.3 Furthermore, when considering the state of the art, the person skilled in the art could not have anticipated that, as demonstrated in the application Example 4, that after incorporation of the TbpB in the OMV of a heterologous meningococcal strain, all variants used for the immunization were able to induce blocking antibodies that were able to inhibit the binding of human transferrin to the meningococcal transferrin receptor, indicating the functional activity of the antibodies. In fact the mixture of Tbps with OMV prepared according to the method of claim 1 was found to confer higher protection than the antigen Tbps alone.
- 3.5.4 However the above effect is only demonstrated with the insertion of the TbpB protein into OMV of *N. meningitidis*, Example 4 and Example 6 involving the incorporation of PorA into OMVs from *Neisseria lactamica* and *Branhamella catarrhalis*. Consequently the subject-matter of claims 2 to 4 and 6 to 12 are considered to involve an inventive step as required by Article 33(3) PCT.
- 3.5.5 The IPEA considers the extrapolation of the method and vaccine of the present application to encompass the incorporation of **any** antigen into **any** bacterial outer membrane vesicle as being purely speculative and not based upon any technical evidence or facts.
- 3.5.6 Consequently the subject-matter of claims 1 and 5 have not been demonstrated as solving the above defined problem and therefore cannot be recognized as involving an inventive step according to Article 33(3) PCT.

Re Item VIII

Certain observations on the international application CLARITY (Art.6 PCT)

INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY (SEPARATE SHEET)

PCT/CU2004/000012

- Claims 1 and 5 encompass **any** antigen incorporated into **any** Gram-negative bacterial OMV, whereas the description and examples actually only involve the incorporation of PorA into OMV from *Neisseria meningitidis*, TbpB and PorA into OMV from *Neisseria meningitidis* and a synthetic peptide containing the variable region 2 derived from the surface loop 4 of class 1 OMP inserted into OMV from *Neisseria meningitidis*.
- 4.1 Therefore all the exemplified antigens are derived from *Neisseria meningitidis* and inserted into the OMV from *Neisseria meningitidis*. There is no technical evidence to indicate that the method would succeed if carried out using any other antigens or any other gram negative bacterial OMV. The general discussion in the description page 7, lines 18 to 21 is not considered sufficient to infer that simply if the method is effective with one member of Gram negative bacteria, that one may assume that the method may be successfully used with any antigen and any Gram negative bacteria.
- The subject-matter of claims 1 and 5 therefore appears to be entirely speculative, not based upon technical facts and not supported by the description according to Article 6 PCT.